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THE PROBLEM OF THERMAL-EXPANSION STRESSES

IN REINFORCED PLASTICS

By P. S. Turner National Bureau of Standards



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MATIONAL ADVISORY COMMITTED FOR AURONAUTICS

ADVANCE RESTRICTED REPORT

THE PROBLEM OF THERMAL-EXPANSION STRESSES

IN REINFORGED PLASTICS

By P. S. Turner

SUMMARY

Failure of adhesive bonds is attributed to boundary stress concentrations. An analysis of the causes of internal-stress concentrations in rigid adhesive layers leads to the conclusion that stress concentrations can be eliminated in many cases by matching the coefficients of thermal expansion of the component parts. A stress-equilibrium formula for calculating the thermal-expansion coefficients of mixtures involves the density, modulus of elasticity, coefficient of thermal expansion, and proportion by weight of the ingredients. Illustrations of the application of the formula derived include lead-antimony and beryllium-aluminum mixtures, phenol-formaldehyde resin and glass-fiber mixtures, and plastic plywoods. The thermal-expansion coefficients of a number of pure and reinforced plastics are reported. Bonds obtained when thermal coefficients are matched are stable over a wide temperature range.

I. MECHANICS OF ADHESIVES

This report describes a method of compounding a plastic or other mixture or compositions to provide a material having a predetermined desired coefficient of thermal expansion. In the preparation of plastics heretofore it has been found practically impossible to produce a satisfactory bond between a plastic and a metal facing or other metal reinforcement. This is due for the most part to the difference in coefficients of expansion of the materials; the plastics have relatively high coefficients of expansion as compared with those of metals. Because of the differential expansion, forces created upon changes in temperature have been such as to prevent a satisfactory bond being obtained between metals and plastics except where plastics of a

flexible rubber-like, or gummy type are used. Even in the bonding of foils of tin, lead, aluminum, etc., to cardboard or paper for containers, gummy or tacky plastics only have been used.

The problem of obtaining bonds between plastics or other adhesives and various materials is considered as being substantially the same problem as that of removing or eliminating concentrated stresses at the boundaries. If internal-stress concentrations can be removed, the full strength of the adhesive can be developed to resist external loads.

Bonds produced by adhesives can be divided into two general classes: the rubbery or yielding bond and the rigid bond. In the first category are found most thermoplastic cements, rubber cements, and combinations of thin rubber layers and cements. Adhesives of this class have been found to provide durable bonds between dissimilar materials at moderate temperatures. The rigid or high modulus bond has generally been found unsatisfactory for such applications. A possible exception is found in the use of cold-setting cements of the phenol-formaldehyde and urea-formaldehyde types. The advantages of the first class over the second disappear at reduced temperatures where the adhesive loses its ability to eliminate stress concentrations by yielding with the dimensional changes of the materials bonded. The rigid bond is superior for many purposes to the yielding bond, if it can be obtained, because it produces a stronger and less-yielding product. For composite structural materials subjected to extreme temperature changes a stable rigid bond is imperative.

In attempting to bond various materials together, it has been found that materials having widely different coefficients of expansion cannot be bonded with any rigid cement available. Certain thermoplastic adhesives produce satisfactory bonds at ordinary temperatures, but fail at low temperatures when the bond becomes rigid and is no longer able to yield to changes in dimension of the materials bonded. Certain rubber cements fall into the classification of bonds which are soft enough to yield with changes in dimensions. These yielding bonds, however, do not produce the rigidity required for structural applications and fail at low temperatures for the same reason that rigid plastic bonds fail at ordinary temperatures; namely, difference in rates of expansion and contraction with changes in temperature.

These failures are caused by stresses resulting from differential expansion or contraction. These forces can be reduced by one or more of the following factors.

Effect of Various Factors on Bond Strengths

An analysie of the factors responsible for the production of stress concentrations in a rigid bond must include the following:

- (1) The thickness of the materials bonded
- (2) The thickness of the adhesive layer
- (3) The modulus of elasticity of all components
- (4) Changes in dimensions with changes in moisture content for all materials
- (5) Changes in temperature to be encountered in service
- (6) Coefficients of linear thermal expansion of all materials

Thickness of materials bonded. - It is evident that if the thickness of the materials bonded approaches zero, stresses set up by any other combination of factors must approach zero also. Accordingly, it has been found possible to bond thin metal foils to other materials. A method of eliminating stresses by this means would, however, have very limited practical application. This stress-reducing factor is utilized in the production of mechanical mixtures of materials of microscopic size or in the form of fine filaments, which are incompatible on a larger scale.

Thickness of adhesive. In bonding two pieces of the same material the stresses set up in the bond as the result of restrained dimensional changes in the bonding material would be proportional to the thickness of the bond. It has long been recognized that thin adhesive layers produce stronger bonds than thicker ones.

Modulus of elasticity. - If the modulus of elasticity of any one of two adjacent materials is zero, the stresses set up between them must be zero. This is essentially

achieved by bonds which yield readily and offer little resistance to static loads whether produced internally or externally.

Moisture content. The problem of bonding materials having different dimensional stability with changes in relative humidity can be reduced, if not eliminated, if the surface of the combination is impervious to moisture. At the time of bonding or at the end of the bonding operation the material or materials sensitive to changes in humidity can be made to contain the normal amount of water present under average conditions. No completely satisfactory solution to this aspect of bonding has been found.

Temperature. - Changes in temperature cannot be eliminated in any practical application. For cold-setting phenol-formaldehyde and urea-formaldehyde cements the effect of temperature changes is minimized by bonding at temperatures close to those encountered in service.

Thermal expansion - From the standpoint of the thermal stresses produced, the problem of bonding two different materials becomes identical with the problem of bonding two pieces of the same material provided the coefficients of thermal expansion of the two materials are the A further reduction is stresses can be achieved if the bonding agent has the came ocefficient of thermal expansion as the materials bonded. A bond between materials having different coefficients of expansion would have to withstand the shear stresses produced by the differential expansion or contraction experienced within the range of temperatures encountered. Bonds can sometimes be maintained if the materials bonded are thin enough to permit the relief of etresses by bending or warping, either of which is undesirable. The bonding strength exhibited by a particular adhesive is the additional load which, added to the internal concentrated stresses, is sufficient to disrupt the material. In some cases this additional load is zero and the materials are said not to bond.

The logical conclusion to be drawn from the foregoing discussion is that stable rigid bonds are possible over.a wide range of temperatures if the thermal-expansion coefficients of the component parts are matched. An example of the results to be expected is furnished by the combination of concrete and steel which individually have practically equal coefficients of thermal expansion and contraction.

It is noted that certain proportions of cement, sand, and stone produce optimum results.

Various methods have been used in other fields to predetermine the thermal-expansion coefficients of mix-tures, such as alloys, glasses, etc. Some of these methods appear to work satisfactorily in limited applications. Notable among these are weighted averages based on the thermal-expansion coefficients of the components and their proportions either by weight or by volume. These methods do not, however, give satisfactory results for plastic compositions.

II. DEVELOPMENT OF A FORMULA FOR CALCULATING COMPFICIENTS OF THERMAL EXPANSION OF MIXTURES

If it is considered that an internal stress system exists in a mixture such that the stresses are nowhere sufficient to disrupt the material, the sum of the internal forces can be equated to zero and an expression for the thermal-expansion coefficient of the mixture is obtained. When small particles or fine filaments are incorporated into a mixture, the small dimensions appear to permit combinations of materials which would be incompatible on a larger scale.

The derivation of the resulting volume thermal ocefficient of a mixture follows:

- a __coefficient of linear thermal expansion
- β coefficient of outloal thermal expansion
- K bulk modulus = 1/bulk compressibility
- d density
- P fraction or percent by weight
- V volume
- AT temperature difference
- S stress

Subscripts:

- n property of nth element
- r resulting property

If it is assumed that each element in the mixture is constrained to change dimensions at the same rate as the aggregate, the stress set-up in the individual particles will be:

$$s_n = (\beta_r - \beta_n) \Delta T V_n K_n$$

since S = strain per unit volume X volume X bulk modulus.
But

$$\Sigma(S_1 + S_2 + \dots S_n) = 0 = (\beta_r - \beta_1) \Delta T (V_1 K_1)$$

$$+ (\beta_r - \beta_2) \Delta T (V_2 K_2)$$

$$+ \dots (\beta_r - \beta_n) \Delta T (V_n K_n)$$

Since

$$\Sigma (\nabla_1 + \nabla_2 + \dots \nabla_n) = \nabla_r, \quad \nabla_n = \frac{P_n d_r \nabla_r}{d_n}$$

which can be substituted for V_n in the above expression. Also, since ΔT , d_r , and $.V_r$ are common factors, they

can be eliminated from each term of the expression. Solving for \$\beta_{\text{the}}\$ the following expression is obtained.

$$\beta_{r} = \frac{\frac{\beta_{1}P_{1}K_{1}}{d_{1}} + \frac{\beta_{2}P_{2}K_{2}}{d_{2}} + \dots + \frac{\beta_{n}P_{n}K_{n}}{d_{n}}}{\frac{P_{1}K_{1}}{d_{1}} + \frac{P_{2}K_{2}}{d_{2}} + \dots + \frac{P_{n}K_{n}}{d_{n}}}$$

Since the coefficient of linear thermal expansion is directly proportional to the cubical coefficient, a can be substituted wherever \$\beta\$ appears with the following result:

$$\alpha_{r} = \frac{\frac{\alpha_{1}P_{1}K_{1}}{d_{1}} + \frac{\alpha_{2}P_{2}K_{2}}{d_{2}} + \dots + \frac{\alpha_{n}P_{n}K_{n}}{d_{n}}}{\frac{P_{1}K_{1}}{d_{1}} + \frac{P_{2}K_{2}}{d_{2}} + \dots + \frac{P_{n}K_{n}}{d_{n}}}$$
(1)

It is apparent by inspection that equation (1) based on stress equilibrium reduces to a percentage by volume calculation if the ingredients have the same bulk moduli. If the ingredients have the same modulus to weight ratios, the calculation amounts to a percentage by weight interpolation.

Equation (1) has been verified with experimental values of several metallic mixtures. The thermal coefficients calculated according to equation (1) for several mixtures of lead and antimony and of beryllium* and aluminum are compared with the measured values in figures 1 and 2.

III. APPLICATIONS TO PLASTIC COMPOSITIONS

In studies of the physical properties of reinforced plastics it has been found that the strongest materials were produced by oriented fibers in thermosetting resins.

^{*}Mixtures of Be and Al apparently form a mixture but no value for the bulk modulus or bulk compressibility was available. From the thermal coefficient, 17.8 × 10⁻⁶/0, of a known mixture containing 32.7 percent Be and 66.3 percent Al, and with reported densities and thermal coefficients of the ingredients, the bulk modulus of Be was calculated from equation (1) to be 15.2 × 10⁶ lb/sq in. Using this value the thermal coefficients of other mixtures of Be and Al were calculated with the results shown in fig. 2 in comparison with the reported values.

The material fabricated in this manner is essentially non-isotropic in all of its properties and has different coefficients of expansion depending on the direction of the fibers in the test specimen. Materials having isotropic properties can be obtained by random distribution or by planned orientation. Solutions for any property must take into consideration the orientation of the filler as well as the percentage composition.

The derivation of the relationships involved follows the same line of reasoning used for the bulk determination. Consider a bar made of filaments of one or more types of material bound together by a resin binder. If this bar is heated, each of the elements would expand a certain amount if not constrained. Since the combination has its own rate of expansion different from that of any of the elements, a certain force would be required to deform each element to the equilibrium length of the composite material. This force in each case would depend on the cross-sectional area of the material, the distance it is deformed from its normal length, and its modulus of elasticity. The external resultant of the sum of these forces must be zero.

Letting A_n equal the area of the element in a plane normal to the axis of the bar with the other symbols representing the other quantities as in the bulk determination (equation (1)), except that Young's modulus (E) replaces the bulk modulus, it is then possible to write:

$$(\alpha_r - \alpha_1) \Delta T (E_1A_1) + (\alpha_r - \alpha_2) \Delta T (E_2A_2)$$

$$+ \cdots (\alpha_r - \alpha_n) \Delta T (E_nA_n) = 0$$

But

$$A_n = \frac{A_r P_n d_r}{d_n}$$

Therefore the expression for the linear thermal coefficient of the mixture under stress-equilibrium conditions is obtained:

$$\alpha_{r} = \frac{\frac{\alpha_{1}P_{1}B_{1}}{d_{1}} + \frac{\alpha_{3}P_{3}E_{3}}{d_{2}} + \dots + \frac{\alpha_{n}P_{n}E_{n}}{d_{n}}}{\frac{P_{n}E_{n}}{d_{1}} + \frac{P_{3}B_{3}}{d_{2}} + \dots + \frac{P_{n}E_{n}}{d_{n}}}$$
(2)

The use of Young's modulus is necessary for oalculating the coefficients of nonisotropic mixtures. The result obtained may be somewhat erroneous because the lateral deformation is neglected.

Application to a Mixture of Polystyrene

and Aluminum Oxide

The applicability of the formula is illustrated by the behavior of brass inserts in a mixture of styrene resin with fused aluminum-oxide powder. Brass inserts in ordinary polystyrene cause the polystyrene to crack bécause of the different coefficients of thermal expansion. The coefficient of linear thermal expansion of polystyrene is approximately 70 x 10⁻⁶/°C, that of brass is approximately 17 x 10⁻⁶/°C.

Fused aluminum oxide was chosen for uge in the mixture because it has a low coefficient of linear thermal expansion (8.7 \times 10⁻⁶/°C) and a high modulus of elasticity compared to its density. Its choice for use with polystyrene was also determined in part by its desirable electrical properties. There was no appreciable change in the excellent electrical resistance of polystyrene when the aluminum-oxide filler was added. The data in table I for the coefficients of linear thermal expansion of mixtures of polystyrene and fused aluminum oxide, calculated according to the stress-equilibrium formula, show that approximately 90 percent of polystyrene and 10 percent of aluminum oxide would be required to match the coefficient of linear thermal expansion of brass. Since the oceffioient of the particular brass used was not accurately known, and since it was thought better to err on the low side, 11 percent of aluminum oxide was used.

In figure 3, curve F represents the values of the coefficient of linear thermal expansion of mixtures of polystyrene and fused aluminum oxide calculated on a percentage by volume basis; ourve G, on a percentage by weight basis; and curve H, by the stress-equilibrium formula (equation (2)). While the thermal coefficient of a mixture of 89 percent of polystyrene and 11 percent of fused aluminum oxide has not been measured, pieces molded with brass inserts show a radical difference between the behavior of pure polystyrene and the filled material. Analysis with polarized light indicates that there are stresses in the pure polystyrene concentrated at the boundaries between the brass and polystyrene, and these are sufficient to rupture the polystyrene. With 11 percent of fused aluminum-oxide filler no such stress concentrations occur and there is no evidence that the filled styrene has fractured. The brass was sufficiently well bonded to the polystyrene mixture to permit sawing and machining of the composite material.

The conventional methods of calculation would yield for this composition a value of 63 × 10⁻⁸/°C for the coefficient of linear thermal expansion on a percentage by weight basis, which would indicate that the material should behave very nearly the same as the pure polystyrene. The behavior of the material, however, indicates that its coefficient is close to that of brass, which value was used to predetermine the composition by the use of the stress-equilibrium formula.

Application to Compositions for Filling

Rivet Depressions

Rivet fillers for aluminum, used to cover depressions caused by riveting, have been tested by exposure to Washington weather for 1 year. An enigma developed when it was found that the composition having the best general weathering characteristics did not stand up the best in this application. The calculated values for coefficients of linear thermal expansion of the various compositions are as follows:

	Material	Calculated coefficient		
(1)	Cellulose nitrate + 25 percent aluminum pigment	44.2 × 10 ⁻⁶ /°C		
 (3)	Cellulose acetate butyrate + 25 percent aluminum pigment	45.2 × 10 ⁻⁶ /°C		
(3)	Vinylite A + 30 percent aluminum pigment	30.6 × 10 ⁻⁶ /°C		

Since the coefficient of linear thermal expansion of aluminum is $24 \times 10^{-6}/^{\circ}$ C, the results are easily explained on the basis that composition (3), which happened to come closest to the coefficient of aluminum, gave the best bond.

Inasmuch as the lowest coefficient obtainable with aluminum pigment is 24 x 10⁻⁶/°C for 100-percent pigment, it becomes necessary to add another ingredient with a coefficient below aluminum to the plastic in order to obtain matched thermal coefficients. The following proportions were calculated for cellulose-plastic compositions to reasonably match aluminum:

Cellulose nitrate or cellulose acetate butyrate

66 percent

Aluminum pigment,

20 percent

Fused aluminum cxide

14 percent

Similarly, the following proportions were calculated for a vinyl-acetate resin composition to match its coefficient of linear thermal expansion with that of aluminum:

Vinylite A resin

74.6 percent

Aluminum pigment

20.0 percent

Fused aluminum exide

5.4 percent

Application to Mixtures of Phenol-Formaldehyde

Regin and Glass Fibers

Approximate values for the properties of glass and phenol-formaldehyde resin required for use in the stress-equilibrium formula are as follows:

Young's medulus of elasticity

For glass
For phenclic resin

11 x 10⁶ 1b/sq in. 1 x 10⁶ 1b/sq in.

Ocefficient of linear thermal expansion

For glass For phenolic resin 7.44 x $10^{-6}/^{0}$ 0 59 x $10^{-6}/^{0}$ 0

Density (g/cc)

For glass 2.56 For phenolic resin 1.30

In order to simplify the calculation of the coefficients of linear thermal expansion of mixtures of these materials, equation (2) can be written in a form consolidating the physical constants:

$$\alpha_{r} = \frac{a_{1}P_{1} + a_{2}P_{2} \cdot \cdot \cdot \cdot + a_{n}P_{n}}{b_{1}P_{1} + b_{2}P_{2} \cdot \cdot \cdot \cdot + b_{n}P_{n}}$$
(3)

where a and b are constants for specific materials for a specific temperature range. The approximate values for these constants for the phenol-formaldehyde resin and glass calculated from the physical constants listed above are:

	Glass		Glass	Phenol-formaldehyde res		
a	•		29.1	45.4		
ъ	X	10 ⁶	3.91	. 77		

In figure 4, ourve I represents a percentage by weight interpolation for the glass-resin mixtures; ourve J is calculated by the stress-equilibrium formula. ured values of the coefficients of linear thermal expansion of two compositions shown in table II are plotted according to the percentage of the resin in the mixture. These samples contained approximately 4 percent of starch which was neglected in calculating curve J. The theoretical coefficients for the compositions containing starch were not calculated because the required physical constants are not known. The experimental points fall much closer to the curve J calculated by the formula than to curve I. The curve on a percentage by weight basis would be similar to curve F in figure 3 and would be still further from predicting the actual result.

The addition of metal faces to reinforced plastics may be desired in order to take advantage of the high-mod-ulus properties of metals, as in an I beam, or to obtain a facing impermeable to moisture. To do this effectively it would be necessary to have matched thermal-expansion coefficients in order to insure satisfactory bonds. To calculate the mixture of glass fiber and phenolic resin required to match aluminum alloy 24S, whose coefficient

of linear thermal expansion is $23 \times 10^{-6}/^{0}$ 0, let α_{r} in equation (3) equal this latter value and solve for the percentage of glass; thus

$$23 \times 10^{-6} = \frac{29.1P_g + 45.4 (100-P_g)}{3.91P_g + 0.77 (100-P_g)} \times 10^{-6}$$

The required percentage of glass is found to be 31.3.

The indicated glass content is seen to be a little higher than was present in the panels prepared with glass-fiber mat, phenolic resin, and starch, whose thermal-expansion coefficients are reported in table IV. These panels were fabricated between aluminum-alloy-24S faces. It was found possible to remove the metal face without seriously damaging it or the core material. The metal faces could not be removed from several other panels, having compositions closer to that calculated, without tearing the metal, shearing the metal at the bond, or rupturing the core.

The starch was added to these compositions in order to absorb the water present in the ingredients or released by the resin during curing. It has been found difficult or impossible to produce plastic compositions with phenolic-thermosetting resins at low pressures unless some water-absorbent material is present. Some of these glassfiber panels were bonded at pressures as low as 50 pounds per square inch.

IV. DETERMINATION OF COMPFICIENTS OF THERMAL EXPANSION OF PLASTICS

In the early stages of the investigation an estimate of the coefficient of thermal expansion of the resinbonded materials was obtained by the following method. Composite strips of one layer of metal and one of a fibrous material impregnated with the resin in varying proportions was pressed flat and cured at 150°C. The strips were then permitted to cool to room temperature. If the composite strip remained straight when cooled, it

was considered that the thermal expansion of the plastic composition matched the metal. As the investigation proceeded, quantitative data for the coefficients of linear expansion of the resins were required.

The equipment used for this purpose consisted of two telescoping fused-quartz tubes, the inner tube being shorter than the outer and resting on the specimen. expansion was measured by a dial gage, graduated to 0.0001 inch, and fastened to the outer quartz tube with the measuring foot resting on the inner quarts tube which in turn rested on the specimen. The temperature of the specimen was varied by immersing the quartz tubes containing the specimen in a water or an oil bath, heated with a hot plate, so that the rate of temperature rise of the bath was approximately 1° C per minute. The temperature of the specimen was measured by means of a potentiometer connected with a copper-constantan thermocouple, which was attached to the center of the specimen. Simultaneous measurements of temperature and extension were obtained by setting the potentiometer at predetermined points and recording the extension when no deflection was produced by opening or olosing the circuit. The ocefficient of linear expansion of the specimen was calculated from the temperature-extension curves. The results thus obtained were corrected for the expansion of the fused-quartz tubes by adding the coefficient of linear thermal expansion of fused quartz, $0.5 \times 10^{-8}/^{\circ}$ d. While this relatively rapid method does not insure a uniform temperature throughout the specimen, it presents a better picture of the changes taking place in the specimen which might be masked by changes in water content and by plastic flow at elevated temperatures, both of which would appreciably affect the results if the time of heating the specimen was prolonged. The stress on the specimen applied by the inner quarts tube and the micrometer foot did not exceed 10 pounds per square inoh.

The specimen used for thermal-coefficient measurements was a bar $6\frac{1}{2}$ to $7\frac{1}{8}$ inches long and about 1/2 inch wide. The thickness was that of the sheet from which the specimen was cut. Specimens for determination of the properties of both pure phenolic resin (Durez 120) and compositions of this resin containing filler were molded in the laboratory at 150° C and 1000-pounds-per-squareinch pressure. The length of the specimen was measured at room temperature to the nearest 0.01 inch with a steel scale. The specimens were rounded on the ends in order

to obtain definite contact between the specimen and the apparatus over a small area. The position of the specimen with respect to the quartz tubes was maintained by washers near the ends of the specimen which came in contact with the sides of the tubes but offered no appreciable resistance to vertical motion of the specimen.

Measurements of the coefficient of linear expansion of an alloy using the method described above are compared in table III with precision measurements on a duplicate sample. The results are not strictly comparable since the coefficient of linear expansion of the sample changes slightly with the heat treatment occurring during measurement. Because the time required to make the measurements differs in the two methods, the amount of annealing that has occurred between measurements is different. The results, however, are essentially the same and indicate that the method employed for the experimental work described in this paper is quite reliable.

Factors Affecting Thermal-Coefficient

Measurements of Plastics

Before considering the actual measurements, some of the factors peculiarly affecting the thermal expansion and related properties of plastics should be discussed. Organic plastic compositions, or water-absorbent materials, in general, in addition to the basic constituents, contain a certain amount of water which apparently acts the same as an additional solid in the composition. A great deal of the variation in the reported values for the linear thermal expansion of wood is undoubtedly related to differences in the water content of the wood at the time of testing. In order to minimize such effects all specimens tested in this investigation were initially conditioned at 25°C and 50-percent relative humidity, which is representative of normal conditions.

The modulus of plastics varies considerably with change in temperature, particularly near the softening point. The range of temperature to which the reported values apply was chosen in each case to avoid large changes in water content and to avoid softening points. Mach of these effects is indicated by breaks in temperature-extension and extension-time curves. Near the softening point of pure plastics the molding strains are

relieved and dimensional changes occur which depend on the method of manufacture. These changes correspond to the dimensional changes encountered on aging. Reinforced or laminated materials also show changes in dimensions involving the relief of the thermal stresses set up by the differential coefficients of expansion of the resin and reinforcing medium. Prolonged heating at elevated temperatures causes shrinkage by the removal of water from the mixture. The relief of molding stresses seems to be permanent, the removal of water reversible.

Results of Thermal-Expansion

Measurements for Plastics

Representative measurements of thermal-expansion coefficients of plastic materials for various temperature ranges are presented in table IV. A typical thermal-expansion curve for pure molded phenolic resin is shown in figure 5.

Micronex or carbon black is frequently added to plastic compositions as a filler. It is therefore interesting to note that the addition of 10 percent of this material to phenol-formaldehyde resin reduced the thermal coefficient of the mixture approximately 20 percent. (See table IV.) From published values for the density and thermal-expansion coefficient of carbon black together with the measured properties of the pure phenolic resin, the bulk modulus of carbon black was calculated to be approximately 3 × 10⁶ pounds per square inch.

Lignin resin is used in a number of applications in conjunction with phenolic resins. Here it is treated as a filler. Two effects are to be noted; first, that the product does not become stable until it has been heated a considerable length of time beyond the time required to cure the phenolic resin; and second, that the lignin initially increases the coefficient of the mixture but after heating finally decreases it by a small amount.

The effect of the walnut-shell flour is similar to that of the lignin resin. The manufacturers of this material state that it contains a large proportion of lignin. One of the uses of this material is as an extender for plywood adhesives.

The behavior of filled phenol-formaldehyde resin containing 25 percent of glass fiber, 71 percent of phenolic regin, and 4 percent of starch, is illustrated by the ourves for mixtures of glass fibers and phenolic resin shown in figure 6. Curves L and 0 are for pure phenolformaldehyde resin and glass fibers, respectively. N was obtained on the first heating cycle. The form of the ourse above 70° C is affected by plastic flow, relief of molding strains, and further ouring of the resin. weight of the inner fused-quartz tube and the pressure of the miorometer foot contribute some of the force producing plastic flow. The shrinkage indicated at 100° 0 on ourve N took place over a period of 5 hours, during which time the temperature was maintained constant. The rate of shrinkage was initially rapid but reached a steady value after about 4 hours. After cooling the sample slowly and then reheating to 100° C the same steady rate of shrinkage was resumed for 4 additional hours, after which the specimen was slowly cooled to room temperature. The total shrinkage amounted to 0.5 percent. Curve M of figure 6 is the thermal-expansion curve for this annealed specimen.

The transparent materials for which data are given in table IV have been used or proposed for use as airplane windshields. The temperature-deflection ourves in each case varied from a smooth curve on the first heating cycle between 50° and 100° C. The curve for the Columbia Resin 39 sample (fig. 7, curve P) showed a decrease in expansion in this range followed by a resumption of the upward trend. No such break occurred in the cooling ourve (fig. 7, ourve Q) or in the curve obtained when the sample was heated a second time (fig. 7, curve R), although the temperature was raised above 110° C. The sample was permanently shortened during the first heating to the extent of 0.5 percent. The specimen at the same time increased in thickness. The curve for methyl-methacrylate resin (fig. 8, curve S) for the range 50° to 100° C showed an increase in expansion followed by a decrease. Again the break did not occur in the occling curve (fig. 8, ourve T). The changes were too small to be measured significantly with a steel scale so a check was run on smaller specimens which were heated in an over at 80° 0 for 18 hours. The indicated change was verified by an inorease in dimensions in the plane of the sheet and a decrease in thickness. This is in agreement with observed dimensional changes on aging. Methyl-methacrylate resin

is one of the best available materials from the standpoint of freedom from crazing, unless subjected to strain, and dimensional stability. The surfaces of both materials appeared to be unaffected by the treatment.

The Pregwood samples listed in table IV were submitted by the Formica Insulation Company and were cut from the same sheets as the test specimens reported upon in NACA Advance Restricted Report of July 1941, entitled "Properties of Reinforced Plastics and Plastic Plywoods." The modulus data reported at that time for parallel-ply material may be used with the measured thermal coefficients to calculate the thermal coefficients of the crossply material. The modulus of the cross-ply material may also be calculated as a composite column using the data for the parallel-ply materials. It is assumed, perhaps erroneously, that the "high" and "low" resin designations indicate definite resin contents. The calculated and observed values for the several properties are shown in table V.

CONCLUSIONS

The implications of the concept and methods of analysis touched briefly in the preceding pages are too numerous to be adequately treated in so short a space. Special applications have been mentioned throughout the paper. Some general conclusions are presented in the following paragraphs.

Stable bonds are possible between large sections if the thermal coefficients of the component parts are matched. The bond under such conditions is not affected by extreme temperature changes.

The method of computing the thermal coefficients of mixtures is applicable to mixtures in general and particularly applicable to problems in reinforced plastics.

Since the magnitude of the stresses distributed throughout the mixture are proportional to the size of the elements, the ultimate strength of the material and its fatigue limits should be raised by reducing particle size. Specifically, the strength of plywood should be increased by the use of thin veneers. At the temperature of fabrication the effect of particle size should reach a minimum.

It is possible to formulate pigmented protective coatings which will have improved adhesion to the coated materials by the use of the thermal-coefficient formula. The thermal-coefficient method described in this paper Offers a rapid and reliable means of Obvaining the basic physical data needed for use in the calculations. With such information the proper combination of materials can be calculated to yield a film matching the base material with respect to coefficient of thermal expansion.

National Bureau of Standards, Washington, D. C.

TABLE I

CALCULATED COEFFICIENTS OF LINEAR THERMAL EXPANSION OF

MIXTURES OF POLYSTYRENE AND FUSED ALUMINUM OXIDE

Compositi	on of mixture	Calculated coefficient of linear thermal expansion		
Fused aluminum oxide (Al ₂ O ₃) (percent)	Polystyrene	By stress equi- librium formula (equation (2))	mixtures on percentage by volume basis	
(percent)	(percent)	(x 10 ⁻⁶ /°C)	(x 10 ⁻⁶ /°C)	
100	o	8.7	8.7	
75	25	9.0	41.7	
50	50	9.6	56.3	
25	75	11.7	64.7	
20	80	12.5	65.9	
9.7	90.3	17	68.2	
5	95	23,2	69.1	
0	100	70	70	

TABLE II

COEFFICIENTS OF LINEAR THERMAL EXPANSION OF MIXTURES

OF GLASS FIBERS, PHENOLIC RESIN, AND STARCH

Compo	osition of mix	Coefficient of linear thermal expansion		
Glass fiber	Phenol- formaldehyde resin (Bakelite XC-11749)	Starch	Calculated by stress equilibrium formula (equation (2))	Measured
(percent)	(percent)	(percent)	$(x 10^{-6})^{\circ}(0)$	(x 10 ⁻⁶ /°C)
100 50 25.2 25	0 50 70.8 75	4	7.44 14.2 	26.0
22.2	73.8 90 100	4	37.4 60	29.9

TABLE III

COMPARISON OF PRECISION METHOD AND RAPID METHOD OF

MEASURING COEFFICIENTS OF LINEAR THERMAL EXPANSION

Tempera	Temperature Average coefficients of linear thermal expansion					
range		Precision methoda		Rapid method		
(°C)		First test Repeat test (x 10 ⁻⁶ /°C) (x 10 ⁻⁶ /°C)		First test (x 10 ⁻⁶ /°C)	Repeat test (x 10-6/00)	
20 to	60	17.7	17.8	17.5	17.2	
60 to	100	18.6	18.3	17.9	18.7	
100 to	150	18,3	18.5	18.4	19.0	
150 to	200		18.7	19.0	19.3	
20 to	100	18.1	18.0	17.7	17.9	
20 to	150	18.2	18.2	18.0	18.3	
20 to	200		18.4	18.3	18.6	
	_					

^aData supplied by Division II-6. The cooperation of P. Hidnert in supplying this information and other data represented graphically in figs. 1 and 2 is gratefully acknowledged.

TABLE IV

MEASURED COEFFICIENTS OF LINEAR THERMAL

EXPANSION FOR VARIOUS PLASTICS

		,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,
		Linear thermal
Material ^a	Range	expansion
		coefficient
	(°C)	$(x 10^{-6}/^{\circ}C)$
Pure phenolic resin (Durez No. 120)	20-100	58.5 - 58.9
and familiary form (addon 110, and, a	20- 50	49.7
	20- 30	
Durez No. 120, 10% Micronex carbon	20-100	
, 20, 50	20-100	
Durez No. 120, 41% reagent-quality		
	20- 50	34.7
aluminum oxide		[
heating)	20- 53	
	53-100	78.8
	20-100	62.8
Durez No. 120, 10% lignin resin		
	20-100	
Durez No. 120, 10% lignin resin	20-100	
(Third heating)	20- 53	-
	53-100	59,9
Durez No. 120, 10% walnut-shell-flour		
filler (First heating)	20- 49	
	49-100	_
* • • • • • • • • • • • • • • • • • • •	20-100	59.2
Durez No. 120, 10% walnut-shell-flour		
filler (Second heating)	20- 49	1
	49-100	
02 V 1 054 D 1 211 T5 22710	20-100	54.1
Glass Mat 25%, Bakelite XC-11749		
phenolic resin 71%, starch 4%	20- 65	00.0
(First heating)	20- 65	1
(Third heating)	20- 65	29.3
atomah 14	20- 70	26.0
Glass Cloth 44%, Durez No. 120 39%,	20- 70	20.0
starch 17%	20- 70	17.8
Columbia resin No. 39	1 20- 10	1,.0
(First heating)	10- 50	103.3
(Second heating)	20- 50	
Dodo	20-100	1
Methyl-methacrylate resin	20- 50	

TABLE IV (Continued)

Material ²	Range	Linear thermal expansion coefficient (x 10 ⁻⁶ /°C)
	20- 50 20- 50	
Pregwood P-1, parallel-ply maple, high resin content Parallel to grain	20- 50 20- 50	
Three-ply direction	20- 50 20- 50	19.8
Three-ply direction	20- 50 20- 50	
cal cross-ply construction, 39.5% Amberlite PR-14	20- 30	15.4

^aPercentage figures are based on the weight of the total composition.

bThese Pregwood materials (P-1 to P-8, inclusive) were erroneously reported to have been prepared with birch veneers in the NACA advance restricted report of July 1941, entitled "Properties of Reinforced Plastics and Plastic Plywoods."

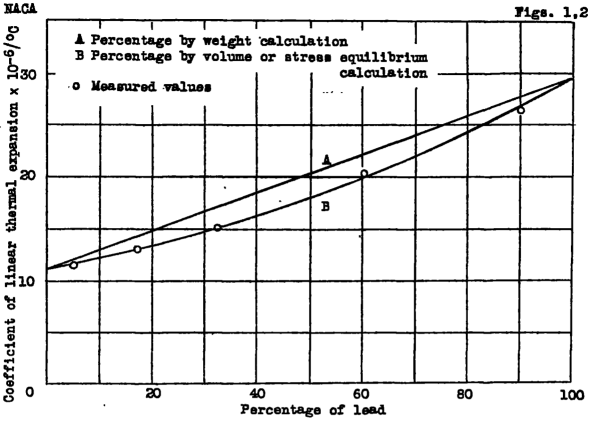
TABLE V

MEASURED AND CALCULATED VALUES FOR COEFFICIENTS OF LINEAR THERMAL EXPANSION

AND MODULUS OF ELASTICITY OF PREGWOODS

	Measured values ^a		Calculated values	
Material	Modulus of elasticity in tension (10 ⁶ lb/sq in.)	Coefficient of linear thermal expansion for the range 20° to 50° C (10°6/°C)	Modulus of elasticity in tension (10 ⁶ lb/sq in.)	Coefficient of linear thermal expansion for the range 20° to 50° C (10° 6/°C)
	I. H	IGH RESIN CONTENT		
	Parallel	-Ply Material (P-1)		
Parallel to grain Perpendicular to grain	3.70 .84	5•9 68•7		
	Cross-Ply	Material, 7 Ply (P-3)		
Parallel to 3-ply direction Do 4-plydo	2.04 2.29	21.0 16.7	2.07 2.47	20 . 5 15.0
	II.	LOW RESIN CONTENT		
	·			
Parallel to grain Perpendicular to grain	3.97 .68	4.7 63.5		
Parallel to 3-ply direction Do 4-plydo	2.17 2.52	19.8 13.5	2 .09 2 . 56	15.6 11.4

The values given for the coefficient of linear thermal expansion are those obtained on the first heating cycle.



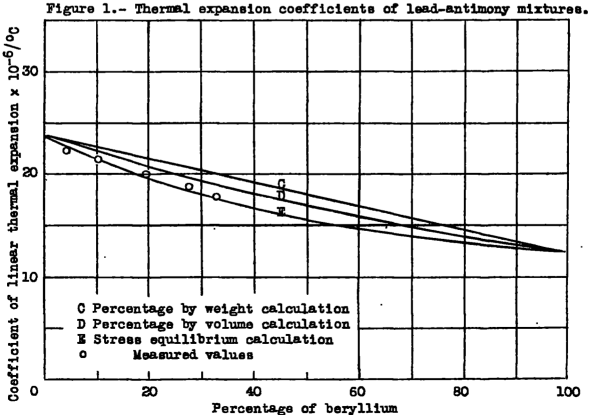


Figure 2 .- Thermal expansion coefficients of beryllium-aluminum alloys.

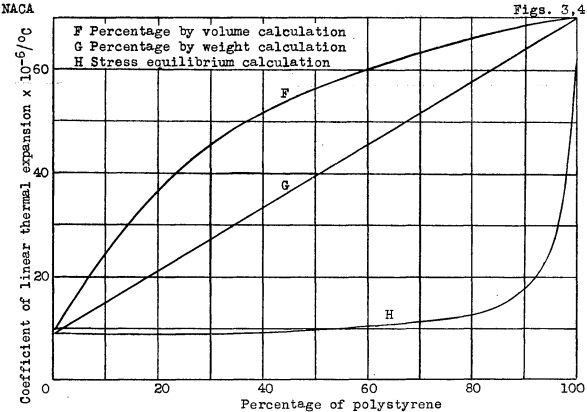


Figure 3.- Thermal expansion coefficients of polystyrene and fused aluminum oxide mixtures.

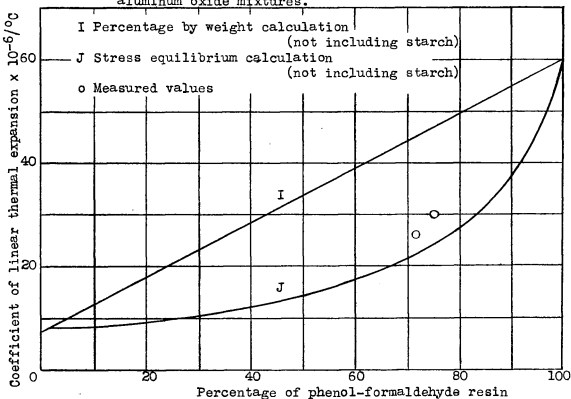


Figure 4.- Thermal expansion coefficients of phenol-formaldehyde resin and glass fiber mixtures containing approximately 4 percent of starch.

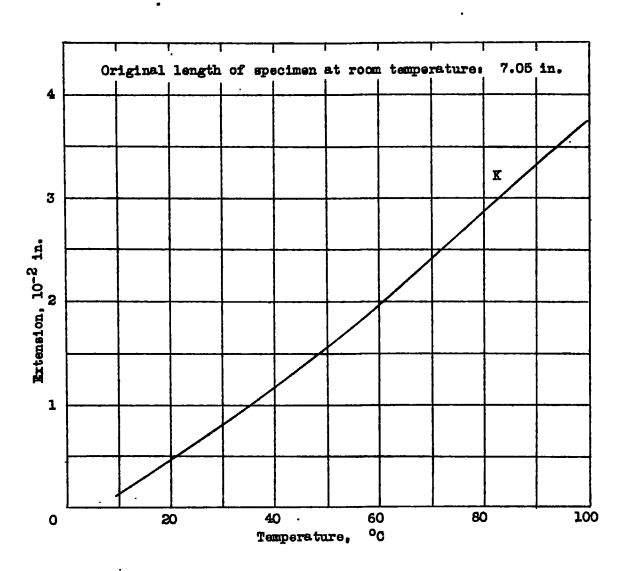


Figure 5.- Thermal expansion curve for pure molded phenolic resin (Dures 120).

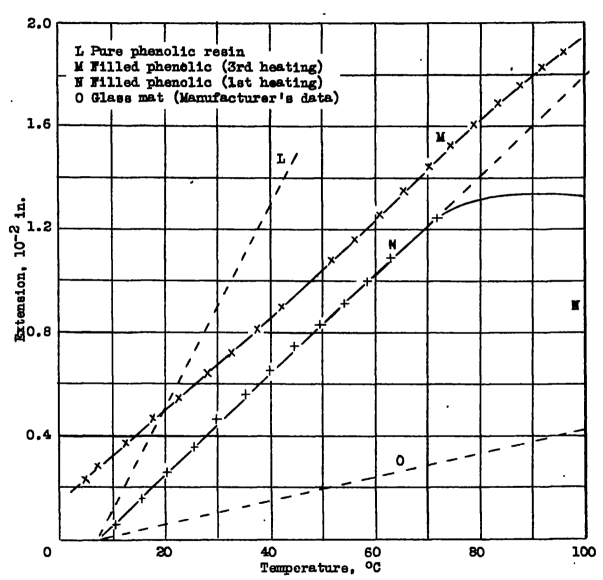
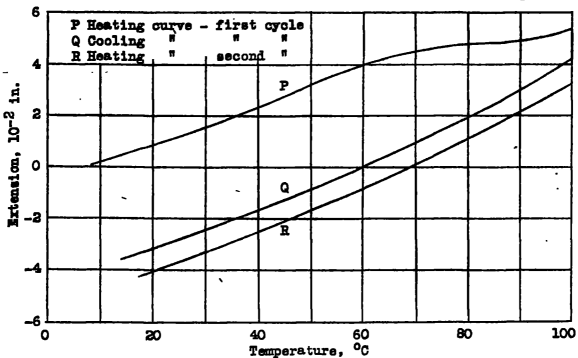
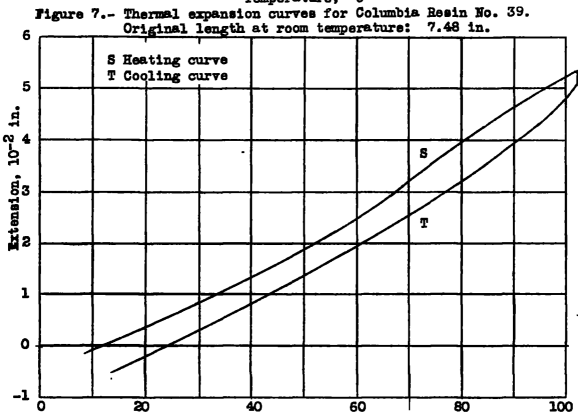


Figure 6.- Thermal expansion curves for phenolic resin and glass fiber mixtures.

Original length at 25° C: 6.57 in. Final length at 25° C: 6.54 in.





Temperature, °C

Figure 8.- Thermal expansion curves for cast methyl methacrylate resin.

Original length at 25° C: 7.04 in.

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    Boundary strees concentrations ere responsible for feilure of edhecive bonds. In-
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 matching the thermal coefficients of component parts. Formule ie given for computing
 thermal coefficiente of mixturee, including leed-entimony, beryllium-aluminum, phenol-
 formaldehyde, gloee-fiber mixturee and plestic plywoode. Bombe obtained by matching
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